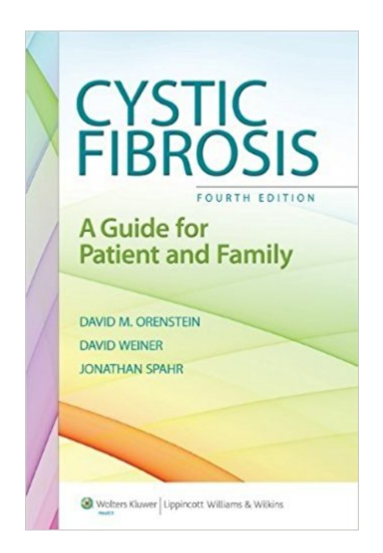


The book was found

Cystic Fibrosis: A Guide For Patient And Family





Synopsis

This one-of-a-kind guide offers easy-to-understand explanations, advice, and management options for patients or parents of patients with cystic fibrosis. The book explains the disease process, outlines the fundamentals of diagnosing and screening, and addresses the challenges of treatment for those living with CF. As one reviewer said, this book â œis the only complete answer book for everyone living with the disease. It is an indispensable resource for families of children with CF, adolescent and adult patients, and physicians, nurses, respiratory therapists, and social workers involved in the care of CF patients.â •

Book Information

Paperback: 416 pages Publisher: LWW; Fourth edition (August 6, 2011) Language: English ISBN-10: 1608317536 ISBN-13: 978-1608317530 Product Dimensions: 1 x 6 x 9 inches Shipping Weight: 1.5 pounds (View shipping rates and policies) Average Customer Review: 5.0 out of 5 stars 14 customer reviews Best Sellers Rank: #189,292 in Books (See Top 100 in Books) #3 in Books > Health, Fitness & Dieting > Children's Health > Cystic Fibrosis #27 in Books > Textbooks > Medicine & Health Sciences > Medicine > Clinical > Gastroenterology #40 in Books > Textbooks > Medicine & Health Sciences > Medicine > Clinical > Pulmonary & Thoracic Medicine

Customer Reviews

This book should be in the home of anyone with CF, a CF family member or knows someone with the genetic disease. The book is written in an easy to understand format. My husband and I walked into our meeting with our new CF team feeling empowered and able to keep up with what the doctors and other team members discussed. I especially like the sections meant to help older kids, teens and adults with CF, there was something in each chapter to help us prepare for the years to come. CF is hard, you want to go screaming for the hills when you first hear the news but pick up this book to help save your sanity!

informative, good format

My daughter was diagnosed in 1997 at 1 month of age. Fortunately for our family we had the best specialist to help diagnose and treat her condition almost from day 1 of her life. And her doctor is none other than Dr. Orenstein himself (or just Dr. O to us). I can speak from personal knowledge that he is without a doubt the most informative doctor I have ever been around. He truly cares about the children in his care and he takes a proactive role in their health and well-being. His book is considered the CF bible by those in the know in the Pediatric Pulmonary field. As you read this picture a man with a bow tie and sandals explaining this to you. It is very easy to read and understand which when it comes to this disease is very important. I am happy to say that my daughter is almost 12 and thanks to Dr. O and his staff is in excellent health. If you know someone with this disease get them this book or better yet get it for yourself as well. Knowledge and understanding of the disease can really help.

Im a teenager and was diagnosed with CF when I was born in 1989. It was recently that I wanted to do some more reading on my disease. I looked through some books and was shocked by what I was reading. As many of you know, the out-of-date books on CF can have some very frightening information, like death and more infections, and etc. It was I discovered this book that I was so much more relaxed about my condition. This book is up-to-date and tells the real facts about this disease. After reading this, my disease doesn't scare me anymore. If you are a teen, like me, I suggest you read thid. If you know anybody with CF, you should read this book or have them read this. Don't let them read those early books on CF with the grim facts...have them read this with the true and helpful facts.

i was given this book as a 'free' gift from a drug company that makescystic fibrosis medication. my son diagnosed was diagnosed at age 5 when he had his pre-k physical. yes...it was very useful after my husband and myself could not find any detailed UPTODATE material on cystic fibrosis...thank goodness more helpful information and websites on cf are out now for newly diagnosed patients and their parents...new parents stay away from grandma's old medical guide on cf you will only find very brief and very grim outdated material...ask your medical team for info or search the web for info on cf

This is a great book. My daughter has CF and this is the book the Doctors gave us. My daughter is treated at one of the top CF centers in the world, so if they use this book only that said a lot to me. I read another before this one and this is by far the best around. Dr O is talking about an update, but

is really busy with research. they hope to do another soon. either way it still has lots of valuable info in it. priceless.

I eagerly await the 3rd edition of this book. As a fully involved CF parent, my 8y/o daughter's care has benefited from my increased knowledge. I can form theories and question her caregivers from a more informed perspective and that is a win-win situation. If you have a child with CF, you should have this book. I call it the CF Bible.(Although encyclopedia would be a more accurate description.) I feel this book should be given to the parents of all newly diagnosed children.

This book is an essential read for: Anyone who has a child with CF, anyone who has a friend with CF, anyone who has a sibling with CF, anyone who has a spouse with CF, anyone who has a parent with CF, anyone who works with people with CF. In short, if you have any contact at all with someone with CF, this book can help educate you. I have had CF for 25 yrs now, and I learned more than a few things.

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